Profound T cell inhibitory activity of hepatic stellate cells (HSCs) in vitro has recently been described in hepatocellular carcinoma (HCC). In the present study, we investigated the immune inhibitory activity of HSCs in vivo in an orthotopic rat HCC model with lung metastasis. Rats (n=24) were randomly sacrificed on days 7, 14, 21 and 28 (n=4 each). Lung tissues were stained with hematoxylin and eosin. Liver sections were stained for immunofluorescence analysis. T-cell apoptosis was detected using double staining for terminal deoxynucleotidyl transferase-mediated dUTP nick-end labeling (TUNEL). Staining revealed marked and continuous accumulation of α-smooth muscle actin with tumor progression after orthotopic tumor implantation in rat liver. T lymphocyte numbers gradually increased following tumor progression, and subset analysis revealed an increase in the distribution of liver CD8+ and CD4+ T cells. Double staining for CD3 and TUNEL demonstrated T-cell apoptosis. Apoptotic T cells were more frequent in the HCC livers compared to the normal livers, and were spatially associated with intratumoral activated HSCs (tHSCs), suggesting a direct interaction. T-cell apoptosis was more frequently induced in the co-cultures of activated splenic T cells (aT)/tHSCs compared to aT/quiescent (q) HSCs or qT/tHSCs. tHSCs were positively correlated with T-cell apoptosis, and the percentage of T-cells undergoing apoptosis was positively correlated with the number of lung metastasis nodules. T-cell apoptosis may be promoted via an interaction with tHSCs, suggesting that tHSCs regulate T cells and contribute to lung metastasis in HCC.

Introduction

The immune microenvironment, biology and prognosis of hepatocellular carcinoma (HCC) have recently been the focus of extensive research. Local cellular immune status has been shown to correlate with invasion, recurrence and metastasis in HCC (1). The liver's unique immunological properties have also aroused considerable interest (2). Liver fibrosis is the result of chronic liver injury, during which hepatic stellate cells (HSCs) proliferate and differentiate into myofibroblasts and express α-smooth muscle actin (α-SMA) (3,4). They also express intercellular adhesion molecule 1 (ICAM-1) and vascular cell adhesion molecule 1 (VCAM-1) and secret chemokines through CD40 expression, and thus, played a role in the regulation of lymphocyte recruitment and migration (5,6). Two recent studies confirmed that T-cell apoptosis can be induced in liver fibrosis and is spatially associated with activated HSCs, suggesting a direct interaction between lymphocytes and HSCs in liver (7,8).

In vitro and in vivo studies have demonstrated that culture-activated HSCs inhibit the function of T lymphocytes through the promotion of T-cell apoptosis (9,10).

Tumor cell line and animals. The rat HCC cell line, Mca-RH7777 (MRH), was obtained from the American Type Culture Collection (ATCC; Rockville, MD, USA). Buffalo rats (6-8 weeks old) were obtained from the Charles River Laboratories (Davis, CA, USA). All animal studies were conducted in accordance with the animal care
policy of the Fudan University and the Animal Research Committee.

Rat tumor model. Intrahepatic tumor implantation was carried out according to Yang et al (12) with minor modifications. MRH cells were injected into the flank of Buffalo rats. HCC tissue was recovered from the transplanted rats 1 month after tumor cell injection, and cut into cubes ~2x1x1 mm³. The rats were anesthetized with ketamine. A small subxiphoid midline incision was made to expose the left lateral lobe of the liver. A small superficial incision was made in the liver and a cube (size, 2x1x1 mm³) of tumor was transplanted. The liver incision was closed with 7-0 suture to avoid any possible early peritoneal seeding of the HCC. Tumor extensions were determined by ultrasound on days 7, 14, 21 and 28 after tumor establishment.

Lung harvesting. Whole lungs of the recipient rats were harvested on days 7, 14, 21 and 28 for quantitation of metastasis as follows. The trachea of the rat was catheterized with a 22-gauge Angiocath (Becton Dickinson Vascular Access, Sandy, UT, USA) and the lungs were insufflated with 15% India ink solution. The lungs were preserved in Fekette's solution for 24 h, followed by 10% formalin. The tumor nodules on the surfaces of the lungs and within the lungs were counted under an operating microscope at x10 magnification (13).

Hematoxylin and eosin (H&E) staining. Cross-sections of liver were checked for the presence of tumor tissue on days 7, 14, 21 and 28 after intrahepatic tumor transplantation. Rats were randomly selected for sacrifice on days 7, 14, 21 and 28 (n=4 each), respectively. Livers were fixed in formalin, embedded in paraffin, and stained with H&E for histological examination.

Immunofluorescence. Liver biopsies taken at all check-points were incubated overnight at room temperature with phosphate-buffered saline (PBS), 10% sucrose and 4% formaldehyde solution. Specimens were frozen at -70°C and 7-μm-thick frozen sections were prepared using a cryostat (Leica CM3000; Meyer Instruments, Houston, TX, USA). Cellular hyperpermeability was achieved using the 0.3% Triton (Sigma, Steinheim, Germany). Bovine serum albumin 1% was used to block non-specific background staining. Following washing with PBS, the slides were incubated with primary antibody to lymphocyte markers and anti-α-SMA rabbit primary monoclonal antibody (mAb) (Roche, Basel, Switzerland) at a dilution of 1:750 for 1 h at room temperature in the dark, then washed three times with PBS. The secondary antibody Cy-5 (BD Biosciences, Franklin Lakes, NJ, USA) was then added, followed by three washes with PBS. In each double-stained set, a single lymphocyte subset was stained using anti-α-SMA. Fluoresceine isothiocyanate (FITC)-conjugated mouse anti-rat CD4 and allophycocyanin (APC)-conjugated mouse anti-rat CD8 cell mAbs (BD Biosciences) were used, respectively. Sections were stacked and covered with Fluoromount-G (Birmingham, AL, USA) to maintain staining and prevent fluorescent bleaching. Sections were then stored at 4°C until analysis by confocal microscopy (14,15).

Terminal deoxynucleotidyl transferase-mediated dUTP nick end labeling (TUNEL) method. For the detection of apoptosis, periodate-lysine-paraformaldehyde-fixed cryosections were stained by the TUNEL technique, using an In Situ Apoptosis Detection kit (Roche), according to the manufacturer's instructions. Sections were incubated at 37°C for 1 h with terminal deoxynucleotidyl transferase in the labeling-safe buffer including FITC, and observed under a confocal laser scanning microscope.

Double staining. After TUNEL staining, cryosections were incubated with Cy5-conjugated anti-rat α-SMA mAb (1:50 dilution; Roche) or phycoerythrin (PE)-conjugated anti-rat CD3 mAb (1:100 dilution; Serotec) overnight at room temperature. They were observed under a confocal laser scanning microscope. FITC images and PE or Cy5 images were entered separately into the computer and superimposed using Adobe Photoshop software (Adobe Systems UK, Uxbridge, UK).

Isolation and culture of HSCs. Rats were anesthetized by intraperitoneal injection of ketamine, 10 mg/100 g body weight. HSCs were prepared from the livers of normal Buffalo rats or HCC tissue, as previously reported (16). In brief, the abdominal cavity of the anesthetized animals was opened to canulate the portal vein. The liver was subsequently perfused at a flow rate of 10 ml/min with Gey's balanced salt solution (GBSS) without Ca²⁺ and Mg²⁺, followed by 100 ml of 0.12% pronase E (Roche) dissolved in GBSS with Ca²⁺ and Mg²⁺ (GBSS) for 10 min. Then 100 ml of 0.1% collagenase (Roche) dissolved in GBSS was recirculated for 20 min. The normal liver and HCC tissue were then excised, dissected and incubated for 30 min at 37°C with continuous shaking, with 0.04% pronase E, 0.05% collagenase and 0.002% DNase I (Sigma) in 100 ml GBSS. After digestion, the cell suspension was passed through a 0.22-μm mesh and centrifuged at 480 x g for 10 min. Subsequently, cells were purified by 8% Nycodenz (Sigma) gradient centrifugation. HSCs were grown in DMEM medium (Gibco-BRL) containing 10% fetal calf serum (FCS) before the experiments were performed. Cell viability determined by trypan blue exclusion was >90%. The purity of quiescent (q) HSCs and tHSCs ranged from 90-95%, as determined by desmin immunostaining. tHSC activation was determined by α-SMA immunostaining. qHSCs and tHSCs were obtained by culture on uncoated plastic dishes for 2 days (17).

Culture of dendritic cells (DCs). Bone marrow cells isolated from F344 rat (Davis, CA, USA) femurs were lysed of red blood cells using red blood cell lysis buffer (Sigma), cultured in RPMI-1640 medium containing 10% v/v heat-inactivated FCS, 20 mmol/l hydroxyethylpiperazine-N-2 ethanesulfonic acid, 2 mmol/l L-glutamine, 0.1 mmol/l non-essential amino acids, 1 mmol/l sodium pyruvate, 20 μmol/l 2-mercaptoethanol, and antibiotics (100 U/ml penicillin and 100 mg/ml streptomycin) (subsequently referred to as complete medium) in the presence of rat recombinant granulocyte-macrophage colony-stimulating factor (4 ng/ml) and interleukin-4 (1,000 U/ml) (both from BD Biosciences). Non-adherent cells were released spontaneously from the proliferating cell clusters, harvested, washed, and resuspended in complete medium, as previously described (18).
Co-culture of HSCs and splenic T cells. Proliferation of nylon-wool-eluted Buffalo rat spleen T cells (2x10⁵/well in 100 µl) was stimulated via allogeneic DCs in a one-way mixed leukocyte reaction. Non-stimulated (quiescent, qT) and DC-stimulated (activated, aT) splenic T cells were added to qHSCs or tHSCs (1x10⁶/well) and further incubated for 48 h. Three rats were used for each group. Apoptotic T cells were identified via double staining with PE-conjugated anti-CD3 mAb and TUNEL. Following surface CD3 staining, cells were fixed in 4% paraformaldehyde and permeabilized with 0.1% Triton X-100 and 0.1% sodium citrate. The TUNEL reaction mixture from the Cell Death Detection kit (BD Biosciences, San Diego, CA, USA) was then added according to the manufacturer's instructions. Cells incubated with the labeling solution in the absence of terminal transferase were used as negative controls. The percentage of TUNEL-positive lymphocytes was then calculated.

Confocal microscopy and image capture. A confocal laser scanning microscope (14900 Superfusion System; Radnoti, Monrovia, CA, USA) was used to analyze the stained sections. Fifty images were collected from each liver using a charge-coupled device camera (Hall 100) and analyzed using Zeiss LSM Image Browser software. Image processing was performed using Adobe Photoshop software (Adobe Systems UK) (19).

Quantitative analysis of immunopositive cells. Livers taken on days 7, 14, 21 and 28 were examined. Four rats were used per group for each time period and 10 sections were prepared per animal. The relative areas of α-SMA-positive cells and numbers of CD4⁺, CD8⁺ and TUNEL⁺ cells were counted in the HCC area, and the total area in one microscopic field, at a magnification of x100. One microscopic field at x100 was 11.4 mm²; a total of 57 mm² of the liver sections were therefore examined per animal. Cell density was calculated and expressed as cell number per mm² (7).

Statistical analysis. All statistical analyses were conducted using SPSS 15.0 software. The data were expressed as the means ± standard deviation (SD). Significant differences were evaluated by unpaired Student's t-tests. Correlations among the density of tHSCs, apoptotic T cells and tumour nodules in the lungs were analyzed using Spearman’s rank correlation. P<0.05 (two-tailed) was considered to indicate a statistically significant result.

Results

Tumor model. Orthotopic HCCs were established with no perioperative complications. All rats (100%) developed tumors during the observation period (Fig. 1).

Metastatic lung nodules. A previous study demonstrated metastatic lung nodules at day 30 in an orthotopic MRH hepatoma model (13). This observation was confirmed in our experiments; 3 out of 6 (50%) animals had significant evidence of tumor nodules by day 21 (Fig. 2D), and 6 out of 6 (100%) by day 28 (Fig. 2E). However, tumor nodules were not obvious at days 7 and 14 (Fig. 2B and C). The number of tumor nodules gradually increased with tumor progression. No tumor nodules were found in healthy rats. Results are representative of 2 different experiments with 4 animals in each time subgroup. Results are presented as the means ± SD, P<0.05. Original magnification, x200.
ranged from 2 to 384 at day 21, (mean, 234.5 nodules per animal), and from 54 to 579 at day 28, (mean, 318.7 nodules per animal). There were no tumor nodules in the normal rats, and 0-2 nodules in HCC rats on days 7 and 14 (means, 0.23 and 0.42 nodules per animal, respectively). No animals had tumor nodules at extrapulmonary sites. These results demonstrate that the number of tumor nodules gradually increased with tumor progression (Fig. 2F).

HSC activation in HCC. The expression of α-SMA (a marker of stellate cell activation) was evaluated in HCC. None of the normal animals showed abnormal α-SMA staining (Fig. 3A), except for the normal light staining. In HCC, however, activated HSCs stained positively with Cy-5 anti-α-SMA and appeared in situ as red cells (Fig. 3B-E). Although α-SMA+ cells might include myofibroblasts and vascular cells as well as activated HSCs, this remains the gold standard for identification of HSCs (14). The α-SMA signal increased markedly with time (Fig. 3B-E). The relative α-SMA area was 0.19±0.12% in normal animals (Fig. 3F), which increased significantly following tumor progression, to 0.79±0.55 at day 7 (P=0.001), 1.81±1.6 at day 14 (P=0.03), 3.02±1.7 at day 21 (P=0.04) and 4.14±1.9 at day 28 (P=0.025).

T-cell infiltration in HCC rat liver. Figs. 4-6 show representative stained samples from normal liver and from HCC livers at 7, 14, 21 and 28 days. Slight T-cell infiltration occurred at day 7 (Fig. 4B) after tumor implantation, and this infiltration increased at days 14 (Fig. 4C), 21 (Fig. 4D) and 28 (Fig. 4E), respectively. There were fewer T lymphocytes in normal rat liver sections compared to those in HCC livers at all time points. The relative total T lymphocyte areas in HCC tissue at each time point are shown in Fig. 4F. The data represent the means ± SD, P<0.05. Results are representative of 2 different experiments with 4 animals in each time subgroup. Original magnification, x200.
livers (Fig. 4A). Immunofluorescence staining demonstrated that the number of CD8⁺ cells per unit square (Fig. 5A2, B2, C2, D2 and E2) gradually increased at each sequential check point. Following tumor progression, CD8⁺ cells were found very close to tHSCs. Cy-5-conjugated α-SMA and APC-conjugated anti-CD8 markers are illustrated with single (A1, B1, C1, D1 and E1, red laser; A2, B2, C2, D2 and E2, blue laser, respectively) and merged stains (A3, B3, C3, D3 and E3, purple cells). Following tumor progression, CD8⁺ cells (blue), were only found attached to tHSCs. There were fewer CD8⁺ cells in normal rats than in HCC rats, and numbers increased significantly, reaching a plateau within 14 days. CD8⁺ cells per unit square peaked at day 21, and decreased at day 28. (A) Normal liver. (B) HCC liver at days 7, (C) 14, (D) 21 and (E) 28. (F) Measured relative CD8⁺ cells at each time point of HCC. Results are representative of 2 different experiments with 4 animals in each time subgroup. Data represent the means ± SD, P<0.05. Original magnification, x200.

Figure 5. Double staining for α-SMA and CD8. Co-localization of CD8⁺ cells and intratumoral activated hepatic stellate cells (tHSCs) in situ suggests cell adhesion. Cells were visualized with confocal laser scanning microscopy. Cy-5-conjugated α-SMA and APC-conjugated anti-CD8 markers are illustrated with single (A1, B1, C1, D1 and E1, red laser; A2, B2, C2, D2 and E2, blue laser, respectively) and merged stains (A3, B3, C3, D3 and E3, purple cells). Following tumor progression, CD8⁺ cells (blue), were only found attached to tHSCs. There were fewer CD8⁺ cells in normal rats than in HCC rats, and numbers increased significantly, reaching a plateau within 14 days. CD8⁺ cells per unit square peaked at day 21, and decreased at day 28. (A) Normal liver. (B) HCC liver at days 7, (C) 14, (D) 21 and (E) 28. (F) Measured relative CD8⁺ cells at each time point of HCC. Results are representative of 2 different experiments with 4 animals in each time subgroup. Data represent the means ± SD, P<0.05. Original magnification, x200.

Figure 6. Double staining for α-SMA and CD4. Co-localization of CD4⁺ cells and intratumoral activated hepatic stellate cells (tHSCs) in situ suggests cell adhesion. Cells were visualized with confocal laser scanning microscopy. Cy-5-conjugated α-SMA and FITC-conjugated anti-CD4 markers are illustrated with single (A1, B1, C1, D1 and E1, red; A2, B2, C2, D2 and E2, green, respectively) and merged stains (A3, B3, C3, D3 and E3, yellow-green cells). Following tumor progression, CD4⁺ cells (green) were only found attached to tHSCs. There were fewer CD4⁺ cells in normal rats, and numbers increased significantly at day 7 in HCC rats. No marked increase in CD4 staining was seen at day 14, but CD4⁺ cell numbers were significantly increased at days 21 and 28. (A) Normal liver. (B) HCC liver at days 7, (C) 14, (D) 21 and (E) 28. (F) Measured relative CD4⁺ cells at each time point of HCC. Results are representative of 2 different experiments with 4 animals in each time subgroup. Data represent the means ± SD, P<0.05. Original magnification, x200.
These results suggest the existence of direct cell to cell attachment. CD8+ cells were always seen adjacent to tHSCs. There were few CD8+ cells in normal animals (Fig. 5F), and the numbers increased significantly, reaching a plateau within 14 days. The number of CD8+ cells per unit square (Fig. 5F) peaked at 21 days, and decreased at 28 days (Fig. 5F). The number of CD4+ cells per unit square, like CD8+ cells, also gradually increased at each sequential...
check point. CD4+ cells first appeared in situ at day 7 (Fig. 5F). Cy-5-conjugated anti-α-SMA (Fig. 6A1, B1, C1, D1 and E1) and FITC-conjugated anti-CD4 markers (Fig. 6A2, B2, C2, D2 and E2) are illustrated with single (red and green) and merged stains (Fig. 6A3, B3, C3, D3 and E3). CD4+ cells were much less abundant than CD8+ cells. Like CD8+ cells, however, the CD4 subsets were also mainly found attached to α-SMA-positive tHSCs (Fig. 6A3, B3, C3, D3 and E3). There were fewer CD4+ cells in normal rats, and they increased significantly at 7 days of HCC (Fig. 6F). No marked increase in CD4 staining was seen at day 14, but CD4+ cell numbers were significantly increased at days 21 and 28 (Fig. 6F).

T-cell apoptosis in HCC. Double staining for CD3 and TUNEL identified some HCC-infiltrating CD3+ T cells that were also positive for TUNEL staining (Fig. 7). Following tumor progression, PE-conjugated anti-CD3- (red; Fig. 7A1, B1, C1, D1 and E1) and TUNEL-stained cells (green; Fig. 7A2, B2, C2, D2 and E2) are illustrated as single and merged stains. Cells stained for both CD3 and TUNEL are yellow-green (Fig. 7B3, C3, D3 and E3). These results suggest the existence of CD3+ cell apoptosis. TUNEL+ cells were rare in normal livers (Fig. 7F), but were significantly increased at days 7 and 14 in HCC livers. The number per unit square of TUNEL+ cells increased and peaked at day 28 (Fig. 7F). Double staining for α-SMA (red) and TUNEL (yellow-green) revealed no obvious α-SMA+/TUNEL+ cells in normal rat livers (Fig. 8A), but double-stained cells were clearly found in HCC livers at each time point (Fig. 8B-E). Direct α-SMA+ HSCs and TUNEL+ cell adhesion gradually increased on days 7, 14, 21 and 28. TUNEL+ cells were closely associated with α-SMA+ HSCs at each time point (Fig. 8A3, B3, C3, D3 and E3). Quantitative analysis demonstrated that the percentage of co-localized α-SMA+ HSCs and TUNEL+ cells per unit square gradually increased in the HCC area at each time point, while there was no appreciable increase in normal rat livers (Fig. 8F).

Induction of T-cell apoptosis in co-culture with HSCs. To confirm the results found in HCC sections, HSCs were isolated from normal and HCC livers and cultured. HSCs from normal livers cultured for 2 days demonstrated quiescent features (qHSCs), with round or star shapes, abundant lipid droplets, and a lack of α-SMA expression. HCC HSCs cultured for 2 days (tHSCs) demonstrated activated features (qHSCs), with round or star shapes, abundant lipid droplets, and a lack of α-SMA expression. HCC HSCs and TUNEL+ cells were closely associated with α-SMA+ HSCs at each time point (Fig. 8A3, B3, C3, D3 and E3). Quantitative analysis demonstrated that the percentage of co-localized α-SMA+ HSCs and TUNEL+ cells per unit square gradually increased in the HCC area at each time point, while there was no appreciable increase in normal rat livers (Fig. 8F).

T-cell apoptosis was positively correlated with metastasis in HCC sections (r=0.861, P<0.05). There was a significant increase in CD3+ T cells that were positive for TUNEL staining (r=0.911, P<0.01) (Fig. 11B). These results suggest the existence of T-cell apoptosis in response to HSCs in HCC sections. TUNEL+ cells were rare in normal livers (Fig. 7F), but were significantly increased at days 7 and 14 in HCC livers. The number per unit square of TUNEL+ cells increased and peaked at day 28 (Fig. 7F). Double staining for α-SMA (red) and TUNEL (yellow-green) revealed no obvious α-SMA+/TUNEL+ cells in normal rat livers (Fig. 8A), but double-stained cells were clearly found in HCC livers at each time point (Fig. 8B-E). Direct α-SMA+ HSCs and TUNEL+ cell adhesion gradually increased on days 7, 14, 21 and 28. TUNEL+ cells were closely associated with α-SMA+ HSCs at each time point (Fig. 8A3, B3, C3, D3 and E3). Quantitative analysis demonstrated that the percentage of co-localized α-SMA+ HSCs and TUNEL+ cells per unit square gradually increased in the HCC area at each time point, while there was no appreciable increase in normal rat livers (Fig. 8F).

Figure 9. Phase-contrast microscopy of co-culture of HSCs and splenic T lymphocytes at 24 h. (A) Quiescent T lymphocytes alone (qT); (B) activated T lymphocytes alone (aT); (C) intratumoral HSCs (tHSCs) and aT cells; (D) HSCs and qT lymphocytes; (E) quiescent HSCs (qHSCs) and aT cells; (F) qHSCs and tHSCs. HSCs were prepared by pre-culture of freshly isolated HSCs from normal and HCC livers for 2 days, respectively. Splenic T lymphocytes were prepared by Ficoll gradient centrifugation, and aT cells and qT cells were obtained by in vitro treatment with rat dendritic cells and culture medium alone, respectively. aT cells clearly adhered to tHSCs in aT/tHSC co-cultures. However, qT or aT cells alone, or in co-cultures of qT cells/tHSCs, aT cells/qHSCs and qT cells/tHSCs, T cells were only slightly associated with HSCs. Results are representative of 3 different experiments. Original magnification, x100.
Considerable advances have been made in terms of our understanding of cancer origins. Genetic and cell-biology studies have indicated that the cooperative activity of parenchymal and mesenchymal cells is central to carcinogenesis and cancer progression (20,21). In the liver, hepatocytes represent the major parenchymal cell type, while the mesenchymal compartment is composed of various cell types, including Kupffer cells and HSCs. HSCs are widely accepted as playing a pivotal role in the regulation of liver function and the development of liver diseases.


tHSCs became increasingly prominent in HCC livers, following HCC progression at each time point, accompanied by marked T-cell apoptosis and lung metastasis. Results are representative of four animal experiments in each time subgroup. Data represent the means ± SD.

<table>
<thead>
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<th>Day</th>
<th>tHSCs (%)</th>
<th>T-cell apoptosis (%)</th>
<th>Metastatic lung nodules</th>
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<td>Day 7 (n=4)</td>
<td>0.87±0.14</td>
<td>19.81±5.33</td>
<td>0.23±0.92</td>
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<td>1.92±0.53</td>
<td>26.32±11.38</td>
<td>0.42±0.18</td>
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<tr>
<td>Day 21 (n=4)</td>
<td>3.01±1.28</td>
<td>52.31±18.21</td>
<td>234.50±92.6</td>
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<tr>
<td>Day 28 (n=4)</td>
<td>4.05±1.97</td>
<td>84.12±29.65</td>
<td>318.70±118.5</td>
</tr>
</tbody>
</table>

Discussion

Considerable advances have been made in terms of our understanding of cancer origins. Genetic and cell-biology studies have indicated that the cooperative activity of parenchymal and mesenchymal cells is central to carcinogenesis and cancer progression (20,21). In the liver, hepatocytes represent the major parenchymal cell type, while the mesenchymal compartment is composed of various cell types, including Kupffer cells and HSCs. HSCs are widely accepted as playing a pivotal role in the regulation of liver function and the development of liver diseases.
role in hepatic tumorigenesis following liver injury (22, 23). HSCs have been previously proposed to exhibit the features of antigen-presenting cells and to stimulate lymphocyte proliferation (24). HSCs transdifferentiate into α-SMA+ myofibroblast-like cells and induce T-cell apoptosis in liver injury (7). T-cell apoptosis induced during inflammation contributes to immune homeostasis in the liver (25, 26). HCC is considered to represent a specific type of liver damage, and activated HSCs can promote the proliferation and metastasis of HCC (27). However, the delicate relationship between the immunological characteristics of tHSCs and HCC metastasis is far from clear.

In the present study, we investigated T-cell apoptosis in HCC in an orthotopic rat HCC model, using cubes of tumor transplanted into rat livers (12). We demonstrated that the role of T cells in HCC appeared to be modulated through direct attachment to tHSCs, emphasized by their cell to cell proximity and surface contact. HCC was confirmed by conventional H&E liver staining, and direct contacts between T lymphocyte subsets and HSCs were manifested in HCC livers. We suggest that lymphocytes migrate into HCC tissue as a result of HCC progression (28, 29), and interact with tHSCs. Confocal imaging revealed significant increases in intrahepatic CD8+ and CD4+ subsets following tumor progression at each time point. These results suggest that all lymphocyte subsets infiltrating into HCC probably interact directly with activated HSCs by adhesion.

T-cell apoptosis was demonstrated by double staining for TUNEL and CD3. Apoptosis of T cells was more common in HCC livers compared to normal livers. T-cell apoptosis was also noted to be spatially associated with tHSCs in HCC, suggesting an inducing role for the latter cells. However, the involvement of other cell types, such as macrophages (30), cytotoxic T cells (31) and tumor cells (32), which are reported to possess apoptosis-inducing functions, could not be excluded. To confirm the inducing role of tHSCs, we therefore, conducted in vitro experiments using co-cultures of αT cells and tHSCs. Apoptosis was increased in these co-cultures, but not in co-cultures of αT cells with qHSCs, or γT cells with tHSCs, indicating that tHSCs are responsible for inducing apoptosis of αT cells. Further studies are planned to investigate the mechanism of T-cell apoptosis with regard to the involvement of cytokine production or cell-receptor expression using this culture system. Because the experiment aimed to demonstrate the requirement for direct contact between lymphocytes and myofibroblast-like cells, we therefore, examined the apoptosis-inducing activity of the culture medium obtained from tHSCs. Supplementation with culture medium from tHSCs did not significantly increase the percentage of T-cell apoptosis, confirming the need for direct contact or close proximity with tHSCs for T-cell apoptosis. These results are in good agreement with the liver-section results. Activation of T cells has been reported to enhance their interaction with fibroblasts via a CD2-dependent adhesion pathway (33), and this interaction serves to increase the survival of αT cells by inhibiting apoptosis (34, 35). The results of the current investigation support this hypothesis; while αT cells cultured without HSCs frequently underwent spontaneous apoptosis, they showed a low frequency of apoptosis in co-culture with qHSCs. Furthermore, apoptosis of T cells was not induced in normal livers containing qHSCs, but was induced in HCC livers containing tHSCs, and in co-culture with tHSCs. The mechanism of T-cell-apoptosis induction by tHSCs is not known. Galectin-1 might be involved in this event, because it is produced by tHSCs but not qHSCs (36), and induces selective elimination of T cells (37). Some data have demonstrated that αT cells use CD44 to undergo apoptosis, and dysregulation of this pathway could lead to increased pathogenesis in a number of diseases, including hepatitis (38). The upregulated expression of B7-H1 (PD-L1, CD274) in activated HSCs can induce T-cell apoptosis (9). It has also been reported that activated HSCs induce transmigration of leukocytes via expression of adhesion molecules such as ICAM-1 and VCAM-1 (5, 39). Taken together, the present model demonstrated prominent T-cell infiltration in HCC and indicated that tHSCs play roles in inducing lymphocyte apoptosis in HCC.

The present study indicated that the number of tHSCs in HCC gradually increased at days 7, 14, 21 and 28, with a corresponding progressive increase in the number of T cells. However, the number of apoptotic T cells also gradually increased, and lung metastasis ultimately developed. Statistical analysis showed that tHSCs in HCC were positively correlated with T-cell apoptosis, and that the percentage of T-cell apoptosis was positively correlated with the number of lung metastasis nodules. The present research, thus, demonstrated tHSC-related T-cell apoptosis, suggesting that THSC immunosuppression in HCC may indirectly promote the growth and metastasis of HCC.

In conclusion, the findings of the present study suggest that T lymphocytes extravasate and accumulate in HCC tissues during tumor progression. They attach directly to tHSCs and are induced to undergo apoptosis through interactions with tHSCs, contributing to lung metastasis of HCC. The present study provides the first evidence for an immunological function for tHSCs in HCC. However, further studies are needed to confirm these results and to identify new targets for inhibiting metastasis of HCC.

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